

Europäisches Patentamt

European Patent Office Office européen des brevets

EP 0 826 714 A2 (11)

(12)

EUROPEAN PATENT APPLICATION

(43) Date of publication: 04.03.1998 Bulletin 1998/10 (51) Int. Cl.⁶: C08G 65/00, C09D 171/02, B01D 12/00

(21) Application number: 97114418.3

(22) Date of filing: 21.08.1997

(84) Designated Contracting States: AT BE CH DE DK ES FI FR GB GR IE IT LI LU MC NL PT SE

(30) Priority: 26.08.1996 IT MI961785

(71) Applicant: Ausimont S.p.A. 20121 Milano (IT)

(72) Inventors: · Strepparola, Ezio Treviglio, Bergamo (IT) Fontana, Simonetta Milano (IT)

 Silvani, Rossella Lentate sul Seveso, Milano (IT)

(74) Representative: Sama, Daniele, Dr. et al Sama Patents, Via G.B. Morgagni, 2 20129 Milano (IT)

(54)Method for the removal of water from surfaces

(57) Method for the water removal from a surface. which comprises covering the surface with a composition having specific weight higher than that of the water. and subsequently removing water from the composition by skimming, such composition comprising a (per)fluoropolyether having molecular weight comprised between 300 and 1500 and a non ionic additive having a (perfluoropolyethereal structure similar to that of the solvent, linked to an hydrogenated part of hydrophilic type. the molecular weight of the (per)fluorinated part is comprised between 400 and 1200 and the ratio by weight between (per)fluorinated part and hydrogenated part is comprised between 1.5 and 3.5.

> SAMA PATENTS 9 MAR 1998

RECEIVEL

Description

The present invention relates to a method for removing water from various surfaces. The problem of the water removal from processed materials is particused in the electronics and fine mechanics field.

One of the most used methods (US-A-4,491,531) consisted in the use of CFC (chlorofluorocarbons) mixtures and of an additive chosen from the following classes: cliamines salts, mono- or diesters of the phosphoric acid or mixtures thereof. Such mixtures allowed the water removal from the material surface by skim-

The banning of the CFC use made it necessary the research of alternative mixtures which, even though 15 maintaining the effectiveness shown by CFC, had a lower environmental impact.

US-A-5,125,978 claims the utilization of mixtures comprising a perfluorinated solvent and a non ionic additive. Such mixtures require an amount of additive not lower than 0.1%, generally higher, and the obtained water removal is not always complete. Moreover the boiling temperature of the solvent, usually around 60°C, makes unavoidable remarkable losses of solvent in the environment.

The Applicant has surprisingly found that mixtures tormed by a (per)fluoropolyether and by extremely low amounts of a non ionic additive result particularly effective in the water removal from various surfaces.

The employment of very low amounts of additive, also of 0.03% by weight, makes it easier the complete removal of the the residues of the additive from the treated surface.

The present invention relates moreover to a method for the water removal from a surface, which method comprises covering the surface with a composition having specific weight higher than that of the water, and subsequently removing water from the composition by skimming, such composition being formed by a (pop-fluoropotyther having molecular weight comprised between 300 and 1500 and by a non ionic additive having a structure such as:

wherein n is such that satisfies the parameter K defined below

with R" H; alkyl C₁₋₃

Y= CF3 or F

The R_I radical of (per)fluoropolyethereal type comprises repeating units randomly distributed along the chain of the polymer chosen from: (CF₂CF₂O), (CFYO) wherein Y is equal to F or CF₃, (C₃F₆O),

(CF₂(CF₂)₂)) wherein z is an integer equal to 2 or 3, (CF₂CF(CR₁)O). (CF(CR₁CP) wherein P, is equal to -CF₃. -C₂F₅. -C₃F₇; CR₁R₅CF₂OF₃O wherein R, and R₅ are equal to or different from each other and are chosen from H, Cl or perfluoroallyl, for instance with 1-4 C atoms.

When the additive is of formula (I), the T terminal of the perfluoropolyethereal radical is chosen from -CF $_3$, -C $_2$ F $_5$, -CICF $_2$ CF(CF $_3$)- CF $_3$ CFCICF $_2$ -.

The number average molecular weight of the (perfluoroethereal part (T-OR₁ or CF₂R₂CF₂) is comprised between 500 and 1200 and the ratio by weight (K) between (perfluorinated part and hydrogenated part is comprised between 1.5 and 3.5. Indeed if the ratio is lower than 1.5 it prevails the contribution of the hydrophile. Hydrogenated part and the additive tends to be water-soluble and therefore extracted from the aqueous phase. If the ratio is higher than 3.5 the additive is uneffective in the water removal. In the case of the formulae (I) and (II) such ratio K corresponds respectively to T-OR₁(CFV)L and CF₂OR₁CF₂CL.

In particular the following fluoropolyethereal R₁ can be mentioned as preferred:

(a) -(CF2CF(CF3)O)a(CFYO)b-

wherein Y is F or CF₃; a and b are integers such that the molecular weight is comprised in the range indicated above:

a/b is comprised between 10 and 100; or the repeating units indicated in (a) can be bound

as follows: -(CF₂CF(CF₃)O)_e(CFYO)_b-CF₂(R'₁)_xCF₂-O-

(CF₂CF(CF₃)O)_a(CFYO)_bwherein R'₁ is a fluoroalkylenic group, for instance from 1 to 4 C:

(b) -(CF2CF2O)c(CF2O)d(CF2(CF2)2O)h-

wherein c, d and h are integers such that the molecular weight is comprised in the range indicated above; c'd is comprised between 0.1 and 10; 1/(c-d) is comprised between 0 and 0.05, z has the value) indicated above, h can also be equal to 0 (c) -(CF_2CF(CF_3)O)_4(CFCO)_1.

wherein Y is F or CF₃; e, f, g are integers such that the molecular weight is comprised in the range indicated above; el(f+g) is comprised between 0.1 and 10, f/g is comprised between 2 and 10,

(d) $\cdot (\tilde{C}F_2O)_i(\tilde{C}F_2O(R_i)O)_i(CF(OR_i)O)_i$ wherein R_i is $\cdot \tilde{C}F_3 \cdot C_2F_5 \cdot C_3F_i$; k_i t are integers such that the molecular weight is comprised in the range indicated above; k_i 4 and j_i+k_i 4 are at least equal to 2. $k_i(l_i)$ 1 is comprised between 0.01 and 1000; l_i 1 is comprised between 0.01 and 100;

(e) -(CF₂(CF₂)₂O)₅wherein s is an integer such as to give the molecu-

wherein's is an integer such as to give the molecular weight indicated above, z has the meaning

already defined;

(f) -(CR₄R₅CF₂CF₂O)_i-

wherein R4 and R5 are equal to or different from each other and are chosen from H. Cl or perfluoroalkyl, for instance with 1-4 C atoms, j' being an integer such that the molecular weight is that indicated above; said unit inside the fluoropolyoxyalkylenic chain being linked each other as follows:

-(CR4R5CF2CF2O)p-R'1-O-

(CR₄R₅CF₂CF₂O)_q-

wherein R' is a fluoroalkylenic group, for instance from 1 to 4 C, p' and q' are integers such that the molecular weight is that indicated above; (g) -(CF(CF3)CF2O);--

j" being an integer such as to give the molecular weight indicated above; said units being linked each other inside the fluoropolyoxyalkylenic chain as follows to have a bivalent radical-

-(CF2CF(CF3)O)a--CF2(R1)xCF2-O-(CF(CF₃)CF₂O)_b-

wherein R', has the meaning indicated above, x is 0 or 1, a' and b' are integers and a'+b' is at least 1 and such that the molecular weight is that indicated above.

These structures comprising the repeating units indicated and the methods for preparing them are described in the patents GB 1,104,482, USP 3.,242,218, USP 3.,665,041, USP 3,715,378, USP 3.665.041, EP 148.482, USP 4.523.039 USP 5,144,092, and as to functional derivatives see USP 3,810,874. All these patents are incorporated herein by reference

The (per)fluoropolyether according to the present invention has number average molecular weight Mn 35 comprised between 300 and 1500, preferably between 400 and 800, and is preferably a perfluoropolyether.

The (per)fluoropolyether has preferably structure of the type:

T-O-R,-T wherein R_f has the meaning indicated above and T' is selected from -CF3, -C2F5, -C3F7; T' is selected from -CF3, -C2F5, -C3F7, -CF2H, -CFHCF2, CF2CF2H

Particularly preferred structures are the following:

$$TO(C_3F_6O)_a$$
- $(CF_2O)_b$ - T " (III)

a" and b" are integers such that the molecular weight is within the range indicated by a"/b" comprised between 1 and 40; T and T are as defined above.

$$TO(C_2F_4O)_p(CF_2O)_qT''$$
 (IV)

p and q are integers such that the molecular weight is within the range indicated by p/q comprised between 0.6 and 1.2; T' and T" are as defined above.

wherein s' is an integer such that the molecular weight is within the range indicated; T' and T" are as defined above.

The extreme effectiveness of these compositions allows the use of amounts of additive generally lower or equal to 0.1% by weight, preferably lower than 0.05%.

For the preparation process of the additives, the above mentioned patents can be utilized, for instance by starting from a monofunctional or bifunctional (per)fluoropolyether, i.e. having -COF terminals, according to USP 3,810,874, incorporated herein by reference.

For instance for the preparation of additives wherein X=CH2O and B=OH one starts from the product having -COF terminals. The -COF group is reduced with metal hydrides to give the alcoholic derivative -CH₂OH which by treatment with 1 mole of ethylene oxide gives the monoaddition product -CH2O-CH2CH2OH. The corresponding tosyl derivative is prepared by reaction with the paratoluenesulphonic acid chloride. The tosyl derivative is then reacted with a large excess of monocomponent polyethylenglycol in the presence of potassium terbutylate. For the other bridge bonds X, the teaching of the above mentioned USP 3.810.874 is followed.

Experimental part

The used (perfluoropolyethereal) solvents are commercially available and differ by number average molecular weight and, consequently, by boiling point and viscosity.

The water removal was measured according to the following methodology:

a drop of distilled water (about 0.1 ml) is deposited on a flat surface of a glass crystallization vessel with base area of about 10 cm2. Some ml of the solution under examination containing 0.03% by weight of additive are then added along the walls until covering the drop and it is noticed what happens after 30". The assigned points correspond to the following cases:

- 1 complete removal
- 2 Residue <10%
- 3 Residue >10%
- 4 No removal

The values 1 and 2 are considered satisfactory.

The following examples are given only for illustrative purposes and are not limitative of the present inven-

EXAMPLE 1

W

moles) are dropped in a more necks jacketed reactor, equipped with thermocouple, mechanical stirrer and reflux, containing 300 g of 50% aqueous solution of NaOH and 200 g (1 mole) of the paratoluensulphonic acid chloride dissolved in 800 ml of CH₂O_E, it is left under strong stirring for 8 hours at room temperature and after having added 11 of water it is left under stirring for further 4 hours.

It is brought to neutrality with hydrochloric acid and the organic phase is separated After distillation of the solvent, 746 g of tosyl derivative are obtained which appears as a limpid liquid, characterized by IR, ¹⁹F NMR and H NMR.

105 g (0.12 moles) of the tosyl derivative prepared above are dropped at 60°C in 4 hours in a more necks 15 jacketed reactor, equipped with thermocouple, mechanical stirrer and reflux, containing 15.6 g (0.14 moles) of potassiumterbutylate and 120 g (0.6 moles) of tetraethylenglycol. Then it is acidified with diluted hydrochloric acid and the organic phase is separated. After a second washing with 150 ml of 5% chlorifidric acid the organic phase brought to drieness results to be constituted by 101 g of $C_{\rm F} C_{\rm F}$

A 0.3% solution of CF₃O(G₂F₆O)₃(CF₂O)_{0,2}CF₂CH₂OCH₂CH₂(CCH₂CCH₂CH₂O)₃OH in CF₃O(C₃F₆O)_{1,7}(CF₂O)_{0,1}CF₃, tested according to 30 the method described above, obtains 1 as evaluation.

ESEMPIO 2

A 0.03% solution of CF₂O

 $C_3F_6O)_3(CF_2O)_{0,2}CF_2CH_2OCH_2CH_2(OCH_2CH_2)_5OH$ in $CF_3O(C_3F_6O)_{1,7}(CF_2O)_{0,1}CF_3$, tested according to the method described above, obtains 1 as evaluation.

EXAMPLE 3 (comparative)

In a more necks jacketed reactor, equipped with thermocouple, mechanical stirrer and reflux containing 15.6 g (0.14 moles) of potassiumterbutylate and 63.6 g (0.6 moles) of diethyleneglycol, 105 g (0.12 moles) of the tosylderivative prepared as indicated in Example 1 are dropped at 60°C in 4 hours. Then it is acidified with diluted hydrochloric acid and the organic phase is separated. After a second washing with 100 ml of 5% hydrochloric acid the organic phase brought to dryness results to be constituted by 93 g of

CF₃O(C₃F₆O)₃(CF₂O)_{0,2}CF₂CH₂OCH₂CH₂(OCH₂CH₂)₂OH, characterized by IR, ¹⁹F NMR and H NMR. The product is characterized by K=4.0.

A 0.03% solution of $\label{eq:cf2} CF_3O(C_3F_6O)_3(CF_2O)_{0,2}CF_2CH_2OCH_2CH_2(OCH_2CH_2)_3OH$

in $CF_3O(C_3F_6O)_{1,7}(CF_2O)_{0,1}CF_3$, tested according to the method described above, obtains 3 as evaluation.

EXAMPLE 4

After addition of 300 ml of water to dissolve the formed potassium iodide, the organic phase, separated and dried under vacuum, is constituted by 53 g of CF₃O(C₃F₆O)₃(CF₂O)_{0.2}CF₂OH₂CCH₂CH₂(OCH₂CH₂C)₃OMe, characterized by IR, ¹⁹F NMR and H NMR. The compound is characterized by K=2.4.

A 0.03% solution of $CF_3O(O_3F_6O)_3(CF_2O)_{0,2}CF_2CH_2OCH_2CH_2(OCH_2CH_2)_{\lambda}OMe \ in \\ CF_3O(O_3F_6O)_1 \ 7(CF_2O)_{0,1}CF_3 \ \ tested \ \ as \ \ described$

above obtains 2 as evaluation. EXAMPLE 5 (comparative)

In a more necks jacketed reactor, equipped with thermocouple, mechanical stirrer and reflux, containing 48.5 g (0.06 moles) of CF₃O(O₃F₆O)₃(CF₂O)_{0.2}CF₂CH₂OCH₂CH₂(OCH₂CH₂O₃OH,

6.72 g (0.06 moles) of potassium terbutylate are added and left under vigorous stirring until the complete disappearance of the solid. Then the terbutylic alcohol developed is removed by distillation, 10 g (0.07 moles) of methyl iodide are dropped and left under stirring for 4 hours.

5

30

product is characterized by K= 3.6. A 0.03% solution of CF₃O(C₃F₆O)₃(CF₂O)_{0,2}CF₂CH₂OCH₂CH₂(OCH₂CH₂)₃OMe in

CF₃O(C₃F₆O)_{1,7}(CF₂O)_{0,1}CF₃ tested as described before obtains 3 as evaluation.

EXAMPLE 6

In a more necks jacketed reactor, equipped with thermocouple, mechanical stirrer and reflux, containing 27 g (0.03 moles) of

CF₃O(C₃F₆O)₃(CF₂O)_{0,2}CF₂CH₂OCH₂CH₂(OCH₂CH₂)_AOH.

3.36 g (0.03 moles) of potassiumterbutylate are added and left under vigorous stirring until the complete disap- 15 peragrace of the solid. Then 5.5 g (0.07 moles) of the chloride of the acetic acid are dropped and left under stirring for 2 hours. After addition of 100 ml of water the organic phase, separated and anhydrified under vacuum is formed by 27.6 g of

 $CF_3O(C_3F_6O)_3(CF_2O)_{0,2}CF_2CH_2OCH_2CH_2(OCH_2CH_2)_4OCOMe,$ characterized by IR. ¹⁹F NMR, and H NMR. The compound is characterized by K=2.2.

A 0.03% solution of CF3O(C3F6O)3(CF2O)0,2CF2CH2OCH2CH2(OCH2CH2 -2)4OCOMe in

 $\text{CF}_3\text{O}(\text{C}_3\text{F}_6\text{O})_{1,7}(\text{CF}_2\text{O})_{0,1}\text{CF}_3$ tested as described before obtains 2 as evaluation.

EXAMPLE 7 (comparative)

In a more necks jacketed reactor, equipped with thermocouple, mechanical stirrer and reflux, containing 35 g (0.05 moles) of

 $CF_O^{\circ}(G_3F_6O)_3(CF_2O)_2CF_2COOCH_3$, 4.3 g (0.07 as more) of ethanolamine are added and are left under vigorous string for 3 hours. After addition of 100 ml of 3% hydrochloric acid, the organic phase, separated and anhydrified under vacuum, is formed by 35.6 g of $CF_3O(G_3F_6O)_3(CF_2O)_0.2CF_2CONHCH_2CH_2OH$, characterized by IR, ^{10}F NMR, and H NMR. The product is characterized by VK=7.3.

A 0.03% solution of CF₃O($_{03}F_{e}$ O)₃(CF₅O)₆,2CF₂CONHCH₂CH₂OH in CF₃O($_{03}F_{e}$ O)₁, $_{7}$ (CF₂O)₆, $_{7}$ CF₃ tested as described 45 before obtains 4 as evaluation.

EXAMPLE 8

A 0.03% solution of $C_3O(C_3F_6O)_3(CF_2O)_0.2CF_2CH_2OCH_2CH_2(OCH_2CH_2)_3OH (having K=2.2) in <math>CF_3O(C_3F_6O)_2.5(CF_3O)_0.1CF_3$ tested as described before obtains 1 as evaluation

EXAMPLE 9 (comparative)

A 0.03% solution of

 $CF_3O(C_3F_6O)_3(CF_2O)_{0,2}CF_2CH_2OCH_2CH_2OH$ (having K=8.6) in

CF₃O(C₃F₆O)_{1,7}(CF₂O)_{0,1}CF₃ tested as described before obtains 4 as evaluation.

EXAMPLE 10 (comparative)

A 0.03% of

CF₃O(C₃F₆O)₃(CF₂O)_{0,2}CF₂CH₂OH (having K=20.8) in 10 CF₃O(C₃F₆O)_{2,5}(CF₂O)_{0,1}CF₃ tested as described before obtains 4 as evaluation

Claims

 Method for the removal of water from a surface, which method comprises covering the surface with a composition having specific weight higher than that of the water, and subsequently removing water from the composition by skimming, such composition comprising a (per)fluoropolyether having molecular weight comprised between 300 and 1500 and a non ionic additive having a structure such as:

 $\label{eq:with L= X-CH2CH2(OCH2CH2)} $$ wherein $X=CH_2O$; $CH_2NR"$; $CONR"$; $$ CH_2OCH_2CH_2NR"$; CH_2OCOCH_2O $$ $$ $$$

wherein n is such that satisfies the parameter K defined below.

B= OH; SH; NHR"; OCH₃; OCOCH₃. with R"= H; alkyl C₁₋₃ Y= CF₃ or F

wherein the R_f radical of (per)fluoropolyethereal type comprises repeating units randomly distributed along the chain of the polymer chosen from: (CF2CF2O), (CFYO) wherein Y is equal to F or CF3, (C3F6O), (CF2(CF2)2O) wherein z is an integer equal to 2 or 3, (CF2CF(ORt)O), (CF(ORt)O) wherein Rr is equal to -CF3, -C2F5, -C3F7; CR4R5CF2CF2O wherein R4 and R5 are equal to or different from each other and are chosen from H, CI or perfluoroalkyl, for instance with 1-4 C atoms, the T terminal of the perfluoropolyethereal radical is chosen from -CF3, -C2F5, -C3F7, CICF2CF(CF3)-, CF3CFCICF2-, CICF2CF2-, CICF2-, the number average molecular weight of the (per)fluoroethereal part (T-OR, or CF2R1CF2) is comprised between 500 and 1200 and the ratio by weight (K) between (per)fluorinated part and hydrogenated part is comprised between 1.5 and 3.5.

55 2. Method according to claim 1 wherein the R_I group comprises the following repeating units:

35

50

55

(a) $+(CF,CF(CF_3)O)_h(CFYO)_b$, wherein Y is F or CF₃; a and b are integers such that the molecular weight is comprised in the range indicated above; ab is comprised between 10 and 100; or the repeating units indicated in (a) can be bound as follows: $-(CF_2CF(CF_3)O)_h(CFYO)_b,CF_2(R^1)_h,CF_2-(CF_2CF(CF_3)O)_h(CFYO)_b$.

wherein R'₁ is a fluoroalkylenic group, for instance from 1 to 4 C;

(b) -(CF₂CF₂O)₃(CF₂OF₂O)₃(OF₃O)₃, wherein c, d and h are integers such that the molecular weight is comprised in the range indicated above; cfd is comprised between 0.1 and 10.7 (Arch) is comprised between 0 and 10.50, z. has the value indicated above, h can also be equal to 0

(c) ·(CF₂CF(CF₃)O)₆(CF₂CF₂O)₇(CFYO)₉ wherein Y is F or CF₃; e, f, g are integers such that the molecular weight is comprised in the average indicated above; e/(f+g) is comprised between 0.1 and 10, f/g is comprised between 2 and 10.

(d) -(CF₂O)₁(CF₂CF₁OR₇)O)₁(CF(OR₇)O)₁, wherein R₇ is -CF₃, -C₂F₃, -C₃F₇; Jk.I are integers such that the molecular weight is comprised in the range indicated above; k+I and j+k+I are at least equal to 2, w(i+I) is comprised between 0.01 and 1000. V_I is comprised between 0.01 and 1000; eq. -(CF₂(CF₃)O₃)-

wherein's is an integer such as to give the molecular weight indicated above, z has the meaning already defined; (f) +(CP_ARsCF₂OF₂O);-

wherein R₄ and R₅ are equal to or different from each other and are chosen from H. Ol or perfluoroalisyl, for instance with 1-4 C atoms, I' being an integer such that the molecular weight is that indicated above; said units inside the fluoropolyoxyaalisylenic chain being linked each other as follows:

-(CR₄R₅CF₂CF₂O)_p--R'_fO-(CR₄R₅CF₂CF₂O)_p-

wherein R_1 is a fluoroalkylenic group, for instance from 1 to 4 C, p' and q' are integers such that the molecular weight is that indicated above:

(g) -(CF(CF₃)CF₂O)_jj" being an integer such as to give the molecular weight indicated above, said units being linked each other inside the fluoropolyoxyalkylenic chain as follows to have a bivalent

-(CF₂CF(CF₃)O)_a--CF₂(R'_t)_xCF₂-O-(CF(CF₃)CF₂O)_b-

radical:

wherein R₁ has the meaning indicated above, x is 0 or 1, a' and b' are integers and a'+b' is at

least 1 and such that the molecular weight is that indicated above.

- Method according to claims 1 or 2 wherein the additive has the structure of type (I) and molecular weight of the perfluorinated part comprised between 500 and 1000.
- Method according to claims 1 or 2 wherein the additive is of type (II) and the molecular weight of the perfluorinated part is comprised between 800 and 1200.
 - Method according to claim 2 wherein the perfluoropolyether belongs to one of the following classes:

$$TO(C_3F_6O)_a$$
" $(CF_2O)_b$ " T " (III)

wherein a" and b" are integers such that the molecular weight is within the range indicated with a "b" comprised between 1 and 40; T' is -CF₃, -C₂F₅, -C₃F₇; T' is equal to T' or -CF₂H, -CF₂CF₂H, -CF+CF₃, as defined above;

$$TO(C_2F_4O)_p(CF_2O)_qT"$$
 (IV)

p and q are integers such that the molecular weight is within the range indicated with p/q comprised between 0.6 and 1.2; T and T are as defined above;

$$TO(C_3F_6O)_s \cdot T^n$$
 (V)

wherein s' is an integer such that the molecular weight is within the indicated range; T'and T" are as defined above.

- Method according to claim 1, wherein the solvent is selected from CF₃O(C₃F₆O)₃, 7(CF₂O)₅, 0CF₃ and CF₃O(C₃F₆O)₂, 5(CF₂O)₅, 1CF₃ and the additive has the formula CF₃O(C₃F₆O)₃(CF₂O)₅ 2CF₂CH₂OCH₂OH₂(OCH₂ CH₃),OH with n=4 or 5.
- 45 7. Method according to claims 1-6 wherein the amount of surfactant in the composition is lower than or equal to 0.1% by weight.
 - Additives according to anyone of the preceding claims.
 - Use of a composition according to claims 1-7 for the removal of water from a surface.